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(54) Thermally polymerized dicyclopentadiene/vinyl aromatic resins and their use

A hydrocarbon resin is prepared by (1) thermally (57) polymerizing a mixture consisting essentially of (a) about 40% to 90% by weight based on total monomers of a cyclic diolefin component comprising at least about 50% by weight dicyclopentadiene, and (b) about 60% to 10% by weight based on total monomers of a specified vinyl aromatic component, preferably alpha-methylstyrene; para-methyl-α-methylstyrene; 2,4-diphenyl-4-methyl-1pentene, or mixtures thereof, and (2) recovering a product having a Ring and Ball softening point of about 70° to about 150°C, having a Mz with a maximum value of about 4000, having a Mw with a maximum value of about 2500, and having a Pd with a maximum value of about 3.0. Hydrogenation of this resin produces light colored, thermally stable products that are useful as tackifiers in _adhesives.

Description

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This invention relates to thermally polymerized copolymers of dicyclopentadiene and vinyl aromatic compounds, a method for making the copolymers, and the hydrogenated products thereof.

Methods are known for thermally polymerizing and copolymerizing dicyclopentadiene (DCPD) feedstocks and for hydrogenating the copolymer product. For example, U.S.P. 4,650,829 discloses a hydrogenated hydrocarbon resin having a softening point lower than 90°C and containing at least 40 weight % of monomer units derived from cyclopentadiene alone and cyclopentadiene plus a comonomer of the class consisting of alkyl-substituted cyclopentadiene, acyclic dienes, vinyl aromatics and mixtures thereof. U.S.P. 5,171,793 discloses the thermal polymerization of a vinyl aromatic component comprising a mixture of styrene and indene and alkylated derivatives thereof, and a cyclodiene at a temperature of 270°C for two hours. The dark colored resin product is mixed with a solvent diluent and hydrogenated to produce a resin having softening points higher than 100°C. U.S.P. 3,040,009 discloses the production of light colored resins by redistilling the bottom fraction of dripolene, a normally liquid mixture of hydrocarbons obtained by high temperature pyrolysis of hydrocarbon gases. The thermally polymerized resin is hydrogenated using a catalyst consisting of metallic nickel distributed on a porous support.

In the past, the molecular weight of cyclopentadiene-based resins was controlled by including codimers of cyclopentadiene with a diene such as methylcyclopentadiene, butadiene, isoprene or piperylene, and also by adding these same linear dienes directly into the polymerization process. In these processes, insoluble waxy polymers were often obtained below a reaction temperature of 250°C. Other methods of softening point and molecular weight control involved not only temperature, but also initial monomer concentration and time in the reactor. Advantages have also been claimed in the final adhesives when indene streams and/or C-9 streams are copolymerized with DCPD streams. However, these C-9 and indene streams are known to contain sulfur, which can create problems in the subsequent hydrogenation process due to early poisoning of the catalyst.

It is known that in a given family of hydrocarbon resins, i.e., all of those based on the same feedstock, as the softening point goes up, molecular weights and polydispersity also tend to increase. It is also known that there is a trend toward higher softening points and molecular weights as reaction times and temperatures are increased during the thermal homopolymerization of dicyclopentadiene. In addition, when vinyl monomers other than alpha-methylstyrenes are used to modify DCPD feedstocks, e.g., styrene and other ring-substituted styrenes such as t-butylstyrene and vinyltoluenes, high molecular weights and polydispersities result.

It would be desirable to provide thermally polymerized resins based on low cost DCPD feedstocks, with relatively low molecular weights and relatively high softening points that do not change substantially with time in the reactor. These resins should preferably readily convert to the corresponding hydrogenated water-white and thermally stable derivatives.

The hydrocarbon resins of this invention consist essentially of a thermally polymerized copolymer made from (1) about 40% to 90% by weight based on total monomers of a cyclic diolefin component comprising at least about 50% by weight dicyclopentadiene, and (2) about 60% to 10% by weight based on total monomers of at least one vinyl aromatic component having the formula

where R_1 is H or a 1-10 carbon linear or branched chain alkyl group at the meta- or para-position and R_2 is a 1-10 carbon linear or branched chain alkyl group or a 2-methyl-2-phenylpropyl group, the resin having a Ring and Ball softening point of about 70°C to about 150°C, having a Mz with a maximum value of about 4000, having a Mw with a maximum value of about 2500, and having a Pd with a maximum value of about 3.0. Alpha-methylstyrene; para-methyl-alpha-methylstyrene; 2,4-diphenyl-4-methyl-1-pentene, or mixtures thereof are amongst the aromatic compounds of use in this invention. The resins are essentially free of halogen, since thermal polymerization avoids the need for Friedel-Crafts catalysts, which produce resins containing organically bound halogen.

Also according to the invention, the thermally polymerized resin can be hydrogenated to produce resins useful as tackfiers for adhesives. The preferred hydrogenated resins are water-white in color and thermally stable. In one embodiment, resins that can be hydrogenated efficiently with minimal poisoning of the hydrogenation catalyst are produced from relatively sulfur-free DCPD feedstocks.

Use of the specified aromatic comonomers makes it possible to control the molecular weight and the softening point during the thermal polymerization. It is thus possible to obtain a resin with a suitably high softening point without rapid increase in molecular weight during the polymerization process.

Monomers suitable for use as the cyclic diolefin component (1) in the manufacture of the hydrocarbon resins of this invention include essentially pure DCPD (at least 95% by weight pure monomer), or mixtures of dicyclopentadiene with codimers of cyclopentadiene and at least one of methylcyclopentadiene, isoprene, butadiene, and piperylene, for example, a mixture containing 50% to 95% or more dicyclopentadiene. Particularly useful DCPD feedstocks are DCPD 101 and/or DCPD 108 available from Lyondell Petrochemical Co., Houston, Texas, U.S.A., which have <1 ppm sulfur. Low sulfur DCPD feedstocks allow more efficient hydrogenation, since there is little sulfur present to poison the catalyst. However, the use of low sulfur feedstocks is not essential for the practice of this invention. Resins made from sulfur-containing DCPD feedstocks can still be hydrogenated, since there are processes that overcome the effect of catalyst poisoning from the sulfur.

The vinyl aromatic monomer comprises at least one compound having the formula

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where R_1 is H, or a 1-10 carbon linear or branched chain alkyl group at the meta- or para-position, e.g., a methyl, ethyl, isopropyl or t-butyl group, and R_2 is a 1-10 carbon linear or branched chain alkyl group or a 2-methyl-2-phenylpropyl group. Alpha-methylstyrene (AMS); para-methyl-alpha-methylstyrene; 2,4-diphenyl-4-methyl-1-pentene (an unsaturated dimer of AMS), or mixtures thereof are of utility. AMS is most preferred. Up to about 15% by weight of total monomers can be a vinyl aromatic monomer other than those specified above, e.g., styrene and styrenes that are ring-substituted with straight chain or branched aliphatic groups, e.g., vinyltoluenes and t-butylstyrene.

The resin product consists essentially of about 40% to 90% by weight based on total monomers of the cyclic diolefin component and about 60% to 10% by weight based on total monomers of the vinyl aromatic component, the total equaling 100%. For the most efficient reaction, the ratio of cyclic diolefin component (1) to vinyl aromatic component (2) is preferably 1:1 or greater. If the amount of vinyl aromatic component exceeds 50%, the yield will decrease, since there will not be enough DCPD to react with all of the vinyl aromatic component. AMS and the other vinyl monomers defined by the formula given above do not homopolymerize. The softening point of the resin product is about 70° to 150°C, preferably about 80° to 140°C.

Reactions can be carried out in solution or neat. When a solvent is used, it is either aliphatic or aromatic. The copolymerization can be carried out in a continuous process, or in a batch mode.

The polymerization is carried out at a temperature of about 210° to 300°C, more preferably about 215° to 280°C, and most preferably about 225° to 275°C. The reaction pressure is self-generated and is typically between 0 and about 150 psig. The reaction time is typically about 10 minutes to about 16 hours at reaction temperature.

The resins of this invention with Ring and Ball (R&B) softening points of $140^{\circ} \pm 5^{\circ}$ C have a maximum Mz of about 4,000, a maximum Mw of about 2,500, and a maximum Pd of about 3.0. Resins with a R&B softening point of $125^{\circ} \pm 5^{\circ}$ C have a maximum Mz of about 3,000, a maximum Mw of about 2,000, and a maximum Pd of about 2.5. Resins with a R&B softening point of $100^{\circ} \pm 5^{\circ}$ C have a maximum Mz of about 2,000, a maximum Mw of about 1,000, and a maximum Pd of about 2.0.

Mz (z average molecular weight) gives an indication of the high molecular weight tail in the resin and has a pronounced effect on the compatibility of the resin in an adhesive base polymer. High Mz is generally undesirable. Mw (weight average molecular weight) is an indication of the average molecular weight of the resin. Mn (number average molecular weight) provides information on the lower molecular weight portions of the resin. Pd (polydispersity) describes the broadness of the molecular weight distribution and is the ratio of Mw/Mn. Mz, Mn, and Mw were determined by size exclusion chromatography using a refractive index detector.

The use of the specified vinyl aromatic monomers provides a reasonably long period of time, i.e., several hours, at selected temperatures during which the hydrocarbon resin can be manufactured while keeping the molecular weight and softening point from substantially increasing with time. It is therefore not necessary to keep the process operating parameters under strict control. For example, when DCPD/AMS at a ratio of 80/20 is thermally reacted in an aliphatic solvent at 60% initial monomer concentration at 240°C, there is a reaction time between 4 and 6 hours, and at 245°C there is a reaction time between 2 and 5 hours during which the softening point remains between 99 and 106°C, Mz and Mw average 1304 and 555 respectively, and Pd is approximately 2.0 or below.

Unlike cationically polymerized products made from similar monomers, thermally polymerized DCPD-based resins retain double bonds that can react further in various ways. The low molecular weight oligomers that are removed from the resin product by distillation before hydrogenation are also reactive. These oligomers can be recycled back into the reaction process with fresh monomer to effectively increase the conversion of monomer to the resin product. Recycle

of the oligomers effectively raises the yield from the typical 50% to 60% to around 90% in the eighth recycle, while still keeping the softening point and molecular weight approximately constant.

Although the unhydrogenated resins tend to be dark in color, they can be used as tackfilers in adhesive systems where initial color is not critical. They can also be used in caulking compounds, sealants, printing inks, waterproofing compositions, and wax compounds. Because the resins retain some reactive unsaturation, these resins can be reacted cationically with other monomers to provide "grafted" terpolymers or "block" copolymer resins.

In order to obtain the highest quality adhesive tackitiers from the copolymers of this invention, it is desirable to hydrogenate the copolymer to lighten the color and remove some or essentially all of the unsaturation, both aliphatic and aromatic, which in turn improves their thermal stability. Halogen and sulfur are essentially not present in preferred hydrocarbon resins of this invention. The feedstocks can be chosen to be essentially free of sulfur, and thermal polymerization avoids the need for Friedel Crafts catalysts such as aluminum chloride that cause chlorine to be organically bound to the resin product. The resins can then be economically hydrogenated with well known hydrogenation catalysts.

Catalysts for the hydrogenation of the copolymer resins of this invention comprise metals selected from Groups VIII, IB, IIB, VIB, and VIIB of the Periodic Table, which are typically used on a support such as aluminosilicate, alumina, carbon, charcoal, or kieselguhr. Group VIII catalysts are preferred, and nickel catalysts are most preferred. Particularly useful catalysts when hydrogenation is carried out in batch autoclaves on sulfur-free resins using slurried catalyst are Ni 5136P and Ni 5256P, available from Engelhard Corporation, Chemical Catalyst Group, Iselin, NJ, U.S.A., and G-96, available from United Catalysts, Louisville, KY, U.S.A. The hydrogenation process can also be continuous, utilizing either continuous stirred reactors or fixed bed catalyst reactors.

The hydrogenation can be carried out neat or in solution. When a solvent is used, aliphatic and/or naphthenic solvents are preferred. The temperature of hydrogenation is about 100°C to 300°C, preferably about 150° to 290°C, and most preferably about 160° to 280°C. The reaction time is typically about 5 minutes to 10 hours, preferably about 10 minutes to 8 hours, and most preferably about 15 minutes to 6 hours.

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The light color hydrogenated resins have good thermal stability that can be improved still further by the addition of small amounts of antioxidants such as, for example, IRGANOX 1010®, a hindered phenol available from Ciba-Geigy, Hawthorne, NY, U.S.A. IRGANOX® 1010 is tetrakis[methylene(3,5-di-tert-butyl-4-hydroxyhydrocinnamate)]methane.

The hydrogenated resins are useful as tackifiers for adhesives, caulking compounds, and sealants; in paints, labels, and tapes; as modifiers, extenders and processing aids for plastics, printing inks, overprint varnishes and other clear coatings; in textile dry sizes, ceramic tile grout, varnishes, waterproofing compositions and wax compounds.

In the following examples, OMSCP means odorless mineral spirits cloud point, which is determined by the following procedure. Ten weight percent resin is mixed in a test tube with odorless mineral spirits. The test tube is then heated until a clear solution is formed. The solution is cooled until turbidity is obtained. The onset of initial turbidity is recorded as the initial cloud point. Cooling of the test tube is continued until visibility is totally obstructed. The final cloud point is recorded at the point of no visibility.

MMAP is the mixed methylcyclohexane aniline cloud point, which is determined using a modified ASTM D-611-82 [1987] procedure. Methylcyclohexane is substituted for the heptane used in the standard test procedure. The procedure uses resin/aniline/methylcyclohexane in a ratio of 1/2/1 (5 g/10 ml/5 ml) and the cloud point is determined by cooling a heated, clear blend of the three components until complete turbidity just occurs. Hydrogenation of the unsaturated portion of a resin, particularly aromatic unsaturation would result in an increase in the MMAP.

R&B softening point is the Ring and Ball softening point, determined according to ASTM E28-67.

To determine the Gardner color (G), 50 weight percent resin is mixed with reagent grade toluene at room temperature until it dissolves. The color of the resin solution is compared to a set of standards on a Gardner Delta Model 211-A Color Comparator, available from BKY Gardner, Inc., Silver Springs, MD, U.S.A. The color values range from 1 to 18, with 18 being the darkest. The notation Gardner 1- (less than Gardner 1) is used to designate a water-white solution.

The Yellowness Index (YID) is obtained by direct readout from a Pacific Scientific Spectrogard™ Color System, Model 96, available from BKY Gardner, Inc., using a measurement cell with a path length of 5.0 cm. The Yellowness Index is useful to distinguish colors lighter than Gardner 1. The closer the YID is to 0, the lighter the color. There is no direct correlation between Gardner colors and the YID, but experience has shown that Gardner 1 is approximately equal to a YID of 40 when both are measured as a 50% solution of resin in toluene and the YID is measured with a measurement cell with a path length of 5.0 cm.

 $UV\alpha$ is a measure of the level of hydrogenation. It is determined in a cyclohexane solution by scanning the spectrum from about 300 nm down to a point where the cyclohexane just begins to absorb. $UV\alpha$ is determined at the point in the scan between 300 nm and the cyclohexane initial absorbance where maximum absorbance occurs. Fully hydrogenated resins exhibit very low $UV\alpha$ values, e.g., about 0.05 to 0.10.

The laboratory thermal polymerization experiments were carried out in conventional stirred one-liter autoclaves available from Parr Instrument Company, Moline, Illinois, U.S.A., equipped with an electronically controlled internal cooling loop and heated externally by an electric heating mantle.

The monomer blend, and solvent when used, was charged to the autoclave and the apparatus was purged with nitrogen before sealing. While stirring, the apparatus was brought to the desired reaction temperature. The heat-up rate

was not critical with times of less than one hour up to twelve hours having been demonstrated. For convenience in the laboratory, quick heat-up was preferred. After the desired time at reaction temperature, the reactor and contents were cooled. The resin was isolated using conventional techniques which included removing solvent and unreacted materials under a nitrogen stream up to 210°C and then low molecular weight oligomers under steam at 225°C. Steam stripping was gentle to help prevent molecular weight increase in the thermally reactive polymer (only 2 ml oil accumulated per each 50 ml of condensed steam water).

The laboratory hydrogenation experiments utilized Parr 1-liter autoclaves. The general procedure was to charge resin, solvent if needed, and catalyst to the autoclave. After flushing to ensure a nitrogen atmosphere, 1400 KPa gauge (200 psig) hydrogen was used for start-up. At 160°C, the hydrogen pressure was adjusted to 6200 KPa gauge (900 psig) and the final reaction temperature was achieved and maintained by selective heating/cooling as needed. When the desired reaction time was complete, the mixture was cooled, vented and flushed with nitrogen. Catalyst was removed by filtration and the hydrogenated products were isolated by distilling off solvent and oligomers. The final distillation was conducted under steam. Other specific conditions are given in the examples.

5 Comparative Example 1

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This example describes the homopolymerization of dicyclopentadiene. No vinyl aromatic monomers are present. A mixture of 360.0 g of DCPD 101 and 240.0 g of recycled aliphatic solvent was heated to the desired reaction temperature in about one hour, using the reaction procedure described above. DCPD 101, available from Lyondell Petrochemical Co., Houston, TX, U.S.A., contains 85% DCPD and <1 ppm sulfur. The reaction time and temperature are given in Table 1, along with the properties of the resin product. The undesirable waxes that formed at reaction temperatures of 240° and 245°C and short reaction times are insoluble in most common solvents.

Table 1

Reaction Temp.	Property	1.0 Hr.	2.0 Hr.	3.0 Hr.	4.0 Hr.	6.0 Hr.							
240°C	Yield (%)	37.4	39.6	59.6	58.0	77.9							
	S.Pt. (°C)	(150)*	(>180)*	112	115	151							
	Mz	483		1106	1234	2944							
	Wax?	Yes	Yes	No	No	No							
245°C	Yield (%)	35.4	44.4	51.4	57.2	74.8							
	S.Pt. (°C)	(160)*	(140)*	116	110	156							
	Mz		424	817	969	2653							
-	Wax?	Yeş	Yes	No	No	No							
250°C	Yield (%)			75.7	63.7								
	S.Pt. (°C)			140	155								
	Mz			2793	2375								
	Wax?			No	No								

^{*}Approximate wax melting points using standard R&B softening point method.

Comparative Example 2

This example describes the copolymerization of dicyclopentadiene with vinyl aromatic monomers other than those claimed in the present invention. The vinyl aromatic component, monomer concentration, reaction temperature and reaction time all varied as shown in Table 2. The monomer concentration is given in parts, with the remainder being

solvent (total = 100).

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-	Gardner Pd Color 2.03 7+						2.03 2.03 2.19 4.99 5.32 5.31
	Mw 617	Mw 617	617 617 636 939	617 656 939 1798	617 656 656 1798 1798	617 656 656 939 1798 3021	656 939 1798 3021 3266 2920
S.P. (°C) Mz	95 1560						
55.2		1.99					
Time (Min.)		45	30	30	45 30 30 480	30 30 480 480	45 30 30 480 45
Temp. (°C)		275	275	275	275 275 275 250	275 275 275 250 275	275 275 275 250 275
Monomer Concentration (parts)		09	99 93	3 3 3	3 3 3	3 3 3 3	3 3 3 3 3
DCPD/VA 84/15	-	85/15	85/15	85/15 75/25 60/40	85/15 75/25 60/40	85/15 75/25 60/40 50/50 40/60	85/15 75/25 60/40 50/50 40/60
Vinyl Aromatic Monomer (VA)		Styrene	Styrene	Styrene Styrene Styrene	Styrene Styrene Styrene Styrene	Styrene Styrene Styrene Styrene	Styrene Styrene Styrene Styrene Styrene Styrene
Sample 1		2	3	2 6 4	2 E 4 2	4 N A	2 6 4 8 9 7

Example 3

This example describes the copolymerization of dicyclopentadiene and alpha-methylstyrene (AMS) at various monomer ratios up to 50/50. The effects of monomer ratio, initial monomer concentration, reaction temperature, and reaction time are shown in Table 3.

The effect of using AMS as the vinylaromatic (VA) comonomer rather than styrene can be seen by comparing Sample 7 of Table 3 (AMS) with Sample 5 of Table 2 (styrene). At a DCPD/VA ratio of 50/50 under otherwise identical conditions except for the VA monomer, even after 8.0 hours of reaction, AMS has limited the Mz to 2292 and Pd to 2.2, while styrene produced an Mz of 18,591 and a Pd of 5.32.

Sample 12 of Table 3 demonstrates that including up to about 15% of other vinylaromatic monomers, based on total weight of monomers, can be accommodated as long as the rest of the vinylaromatic component is AMS. It is therefore

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possible to make terpolymers with controlled molecular weights.

5		Gardner Color	91	8+	10+	11	12-	14+	18	16+	. 12+	12-	> 18	80
		8	1.96	1.99	2.20	1.71	1.74	1.92	2.20	2.71	1.88	1.9	2.78	2.17
10		M *	299	516	814	524	199	735	1084	1420	645	693	1497	288
15		Mz	1404	1362	1919	1015	1084	1511	2292	3402	1327	1462	3541	2193
		S.R. (°C)	105	107	118	18	8	8	ē	91	2	દ્વ	9	25
20		Yield (%)	37.5	60.4	66.7	38.9	40.5	49.7	64.2	63.0	50.9	54.4	73.5	62.5
25	e 3	Reaction Time (Min.)	45	300	300	240	240	240	480	180	240	240	240	15
30	Table 3	Reaction Temp.	275	240	250	225	237	246	250	275	753	230	250	275
35		Monomer Concentration	99	8	09	95	09	09	80	· 09	80	100	001	09
40		DCPD/VA	85/15	85/15	80/20	99/20	50/50	20/20	20/20	80/50	50/50	80/50	. 05/05	70/15/15
45		Vinyl Aromatic Monomer (VA)	AMS	AMS/Styrene										
50		Sample	-	2	6	4	~	۰	,	8	6	2	=	12

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Example 4

This example describes the operating conditions required to prepare a dicyclopentadiene/alpha-methylstyrene (AMS) copolymer at a specified reaction temperature where the molecular weight and softening point are substantially unchanged.

Dicyclopentadiene 101 (288.0 g), 72.0 g of AMS, and 240.0 g of recycle solvent for each reaction were contained in an autoclave and brought to 170°C quickly. DCPD 101 contains 85% DCPD and <1 ppm sulfur and is available from Lyondell Petrochemical Company, Houston, TX, U.S.A. In the next hour the reaction temperature was brought to the desired value as specified in Table 4. After the specified reaction time at temperature, the reaction products were cooled to 170°C in 35 minutes, poured into stripping flasks and finished, solvent distilled under nitrogen up to 210°C, and finally

treated with steam at 225°C.

Table 4

5	Reaction Temp.	Property	1.0 Hr.	2.0 Hr.	3.0 Hr.	4.0 Hr.	5.0 Hr.	6.0 Hr.	7.0 Hr.	8.0 Hr.
Ì	235°C	Yield (%)								66.8
-		S.Pt. (°C)	1							105
		OMSCP (°C)				-				55/-38
10		MMAP (°C)								43/37
ļ		Color								10-
1		Mz `								1700
15		Mw								715
		Pd								2.08
	240°C	Yield (%)			45.3	50.3	60.7	61.8		62.3
		S.Pt. (°C)			(135)*	99	101	106		111
20		OMSCP (°C)			125/-25	135/-30	52/<-50	87/<-50		56/-50
		MMAP (°C)			115/30	102/33	47 <i>/</i> 34	40/37	:	39/37
		Color			7+	7+	9+	10-		10
25		Mz			834	1063	1290	1423		1689
		Mw			393	490	574	641		682
		Pd			1.64	1.76	1.92	1.95		2.12
30	245°C	Yield (%)	33.1	44.5	49.7	58.4	59.3	62.3	69	66.3
		S.Pt. (°C)	(141)*	101	101	104	105	112	129	127
		OMSCP (°C)	135/-30	136/-40	59/<-50	40/-42	61/<-50	44/-45	68/45	45/26
		MMAP (°C)	115/35	116/30	65/36	40/37	38/36	40/37	46/38	43/40
35		Color	7+	7+	8+	9+	9+	10	12	11
		Mz	963	970	1243	1553	1589	1695	2432	2188
		Mw	423	429	498	624	632	732	1030	928
40		Pd	1.67	1.70	1.90	2.00	2.02	2.10	2.43	2.32
	250°C	Yield (%)			54.4	62.1	66.7	67.5		i 1
		S.Pt. (°C)	1		99	117	118	124		
		OMSCP (°C)			32/<-50	15/-32	18/-3	30/2		
45		MMAP (°C)			38/35	47/36	46/40	41/40		-
		Color			9-	10-	10+	- 11-		
		Mz			1189	1666	1919	2086		
50		Mw			536	710	814	871		
		Pd			1.83	2.08	2.28	2.27		

^{*}Approximate wax melting points using standard R&B softening point method.

Example 5

This example describes the copolymerization of dicyclopentadiene and alpha-methylstyrene in which low molecular weight oligomers are recycled to increase the conversion of monomers to resin.

A mixture of 306.0 g of DCPD 101, 54.0 g of AMS, and 240.0 g of recycled aliphatic solvent was brought quickly to 170°C in a stirred autoclave. DCPD 101 is available from Lyondell Petrochemical Company, Houston, TX, U.S.A., and contains 85% DCPD and <1 ppm sulfur. From this point the reactor temperature was raised to 275°C in 20 minutes. After 45.0 minutes at 175°C, the reactor was quickly cooled to 170°C in about five minutes. After transferring the polymerized resin solution to a glass stripping apparatus, the resin was isolated by heating, first under nitrogen and then under steam to give the "0" cycle resin. All the volatiles that were removed from the resin were combined to create a "recycle pool". Recycle #1 was prepared by mixing 240.0 g of volatiles from the original recycle pool with 306.0 g of fresh DCPD 101 and 54 g of AMS and repeating the initial reaction process steps exactly except that the time at 275°C was shortened to 15 minutes. The resin was isolated and the volatile materials were again added to the recycle pool. Seven more recycle experiments were conducted in the same manner with data from selected cycles listed below:

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			Resin Properties											
Recycle No.	Volatiles Yield (% Total Charge)	Yield (% of Mono- mer Charge)	S.Pt. (°C)	Gardner Color	Mz	Mw	Pd							
0	55.6	37.5	105	10-	1404	599	1.96							
1	57.6	57.8	96	9+	1335	550	1.92							
2	43.4	67.7	102	9+	1403	590	1.96							
4	44.6	81.2	98	11-	1636	654	2.11							
8	39.0	89.9	103	12+	2108	892	2.34							

Table 5

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Overall conversion of monomers to resin will be at least 80 to 90% when utilizing recycle volatiles at the 40% level as part of the reactor charge and Mz will be close to 2000. Adjustments in molecular weight and yield can be effected by changes in percent of recycle volatiles utilized, polymerization temperatures, and reaction time.

35 Example 6

This example illustrates the use of DCPD 108 (95% purity DCPD) as the cyclopentadiene source and also shows neat polymerization followed by solution hydrogenation without isolating the desired base resin from the polymerization reaction mixture. DCPD 108 is available from Lyondell Petrochemical Company, Houston, TX, U.S.A., and contains 95% DCPD and <1 ppm sulfur.

Mixtures containing 250 g of DCPD 108 and 250 g of AMS were placed in each of the two one-liter autoclaves. Each mixture was reacted at 250°C for four hours after 3.4 hour heat-up times. After cooling below 150°, the autoclave contents were combined and mixed with 333 g of aliphatic solvent. Ni 5136P (3.13 g) available from Engelhard Corporation, Chemical Catalyst Group, Iselin, NJ, U.S.A. was added to 500 g of this combined resin solution in a 1-liter autoclave at room temperature. During subsequent heat-up, full hydrogen pressure was applied at 160°C to prevent any increase in molecular weight in the base resin. Hydrogenation was carried out for 5.0 hours at 265°C and 900 psig hydrogen pressure. After cooling, purging with nitrogen, and filtering to remove catalyst, a water white product with the following characteristics was obtained: yield of 67.7% (proportionally based on original monomer charge); s.pt. 111°C; MMAP 78°C; color G-1- and YID 11.5 (both measured at 50% by weight in toluene with YID determined in a cell of 5.0 cm path length); Mz 1647; Mw 843; Mn 424, and Pd 1.99.

Example 7

This example illustrates hydrogenation of DCPD/AMS resins at an initial ratio of 80/20.

DCPD 101 was used in all of the experiments listed in Table 6. DCPD 101 is available from Lyondell Petrochemical Company, Houston, TX, U.S.A., and contains 85% DCPD and <1 ppm sulfur. AMS was the vinyl aromatic monomer, and all polymerization reactions were run at 60% monomer in solvent (initial concentration). All other process variables for the hydrogenation are given in Table 6.

Samples 1 and 2 were thermally polymerized in the laboratory using 245°C reaction temperature (4.0 and 5.0 hour samples in Table 4). The isolated resins were then redissolved in aliphatic solvent and hydrogenated under the conditions described in Table 6. Other process conditions not specified in Table 6 are the same as in Example 6. Note the apparent equivalence of Ni 5136P (Sample 1) and Ni 5256P (Sample 2) in these examples. Ni 5136P and Ni 5256P are available from Engelhard Corporation, Chemical Catalyst Group, Iselin, NJ, U.S.A. In the table, G = Gardner color.

Base resin samples 3-6 were prepared in a 50 gallon pilot plant thermal reactor using process conditions similar to those used for producing resins in the laboratory. Solvents and oligomers were removed under vacuum at temperatures of 180° to 200°C. Hydrogenation of the samples was carried out at three different temperatures, three catalyst levels and two different times to evaluate the effect of these variables. Thermal color stabilities are quite good for these types of hydrogenated DCPD-based resins.

15 20 25		4 5. 6		103 103 103	25/12	43/40 4	<u></u> ≛	2191	618	348 348 348 2.35 2.35 2.35			165,0*** 165,0*** 165,0***	N5136P N	00:1	335.0 3	3.0	222	0079	(900 psig) (900 psig) (900 psig)	8:001	112** 109** 103**	-25/<50 -2	01/91	14/17.03	0.30 0.30	7 4+	1 7721	25	198	2.13 2.16 2.16	0.192	0.58			
30	Table 6	3		103	25/12	43/40	±	2191	618	23.5			165.0***	N5136P	1.50	335.0	4.0	052	9700	(900 psig)	95.9	**601	-30/<-50	25/28	1-6-01	0.30	,	7	740	3.5	- S	0.147	0.70			
35		2		105	05->/69	38/36	ま	1589	632	313			165.0	NiS2SGP	05:1	335.0	4.0	જ	6200	(900 psig)	96.2	117	13/<-50	74/70	1-/8.7	0.15	4	1331	100	310	5 5	0000	1.13			
40		1		101	40/-40	40/37	ま	1553	624	312			165.0	N5136P	S .1	335.0	4.0	250	9000	(900 psig)	986	115	-12/<-50	13/1	1-/10.0	0.15	3	۲,	705	3 2	96	0.095	1.07			
45		Sample	Base Resin	(C)	OMSCP (C)	MMAP (°C)	Color (Gardner)					Hydrogenated Resin	W Resin (0)	9	% Catalyst	Aliphatic Solvent	Time (Hr.)	Temp. (°C)	Hydrogen	Pressure (KPa)	-	S.P. (°C)	OMSCP (°C)	MIMAP (°C)	Color (GYID*)	% Inganox 1010	24 Hr. Heat Stab.	METADOR CONCE)				-	Brounine No.	en Chi coll	**Stripped to S.Pt.	***Stripped pilot plant resin.
50			Base	A S	. 3 6	MA	3	¥	Ž	₹ 3	:	圣	3	Catalvs	3	Aisi	Ĭ,	Tem	Hydr	Æ	Y.ek	SP	Sec	Ž	3	% %	24 H	2	ž ž	1	2 2	1170	Bron	3	- S	S***

Example 8

This example illustrates copolymerization of dicyclopentadiene and alpha-methylstyrene using Shell DCPD (<5 ppm sulfur).

A blend of 288.0 g of 75% DCPD (available from Shell Chemical Company, Houston, Texas, U.S.A.), 72.0 g of AMS and 240.0 g of aliphatic solvent was added to a 1-liter Parr Autoclave, flushed with nitrogen and brought to 170°C over a period of 30 minutes. The temperature was increased to 245°C over a period of one hour and maintained at 245°C for an additional 4.0 hours. The resin was isolated by distilling solvent and unreacted materials up to 210°C under nitrogen. Steam stripping at 225°C removed low molecular weight oligomers, leaving a 57.9% yield of copolymer resin with the following physical characteristics: s.pt. 106°C; color G-9+; OMSCP 65/-42°C; MMAP 65/36; Mz 1267; Mw 535; Pd 1.96. Note that this resin has physical properties that are almost identical to a DCPD/AMS 80/20 resin made under the same conditions using DCPD 101 from Lyondell Petrochemical Company, Houston, TX, U.S.A. (see Table 4).

Example 9

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This example illustrates copolymerization of dicyclopentadiene and alpha-methylstyrene using a sulfur-containing feedstock (85% DCPD available from Petroplast Chemical Corp., White Plains, NY, U.S.A.) followed by poisoning attributable to 0.15% sulfur in the Petroplast DCPD (Sample 2). Poisoning is indicated by a lower level of hydrogenation and a darker hydrogenated resin color. A comparison is made with a resin (Sample 1 in Table 7) made under identical conditions using DCPD 101, available from Lyondell Petrochemical Company, Houston, TX, U.S.A.

The mixtures described above were each polymerized in one liter autoclaves using identical conditions, as shown in Table 7. The resulting resin solutions were cooled to room temperature and the autoclaves were opened. Ni 5136P was added to each mixture, the autoclaves were again sealed, flushed with nitrogen and identical hydrogenations were carried out as shown. Ni 5136P is available from Engelhard Corporation, Chemical Catalyst Group, Iselin, NJ, U.S.A. At the end of the hydrogenation cycles, the products were cooled to 170°C, filtered to remove catalyst and finished in normal fashion, finally steaming at 225°C. The evidence that Sample 2, made from the sulfur-containing feed, has undergone less hydrogenation than the clean feed is compelling: lower MMAP, higher UVa and bromine number all show that Sample 2 is less aliphatic than Sample 1. There is also a striking difference in the colors of the two products, with the

sulfur-containing feed resulting in color reduction only to a Gardner 3+ level, which is far from water-white.

Table 7

Sample	1	2
Lyondell DCPD 101 (<1 ppm S)	288.0 g	
Petroplast DCPD (1500 ppm S)(g)		288.0
AMS (g)	72.0	72.0
Aliphatic solvent (g)	240.0	240.0
Polymerization		
Temp. (°C)	245	245
Time (hr.)	4.0	4.0
Hydrogenation		
Ni 5136P (g)	3.00	3.00
Time (hr.)	3.0	3.0
Temp. (°C)	225	225
Resin Yield (%)	53.3	60.8
S.Pt. (°C)	96	108
MMAP (°C)	74/66	66/62
Gardner color (50% toluene)	1-	3+
YID (2 cm Cell)	25.4	49.0
UVα	0.067	0.218
Bromine number	0.41	1.01
Molecular weight		
Mz	1081	1231
Mw	485	540
Mn	270	286
Pd	1.79	1.89

It is not intended that the examples given here should be construed to limit the invention, but rather they are submitted to illustrate some of the specific embodiments of the invention. Various modifications and variations of the present invention can be made without departing from the scope of the appended claims.

45 Claims

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 A hydrocarbon resin consisting essentially of a thermally polymerized copolymer made from (1) about 40% to 90% by weight based on total monomers of a cyclic diolefin component comprising at least about 50% by weight dicyclopentadiene, and (2) about 60% to 10% by weight based on total monomers of at least one vinyl aromatic component having the formula

where R_1 is H or a 1-10 carbon linear or branched chain alkyl group at the meta- or para-position and R_2 is a 1-10

carbon linear or branched chain alkyl group or a 2-methyl-2-phenylpropyl group, the resin having a Ring and Ball softening point of about 70° to about 150°C, having a Mz with a maximum value of about 4000, having a Mw with a maximum value of about 2500, and having a Pd with a maximum value of about 3.0.

- 5 2. The resin of claim 1, wherein the cyclic diolefin component also comprises codimers of cyclopentadiene with at least one compound selected from the group consisting of methylcyclopentadiene, isoprene, butadiene, and piperylene.
- 3. The resin of claim 1, wherein the vinyl aromatic component also comprises up to 15% by weight, based on the total weight of monomers, of a compound selected from the group consisting of styrene and styrenes that are ring-substituted with straight chain or branched aliphatic groups.
 - 4. The resin of claim 1, wherein the ratio of component (1) to component (2) is 1:1 or greater.
- 15 5. The resin of claim 1, wherein the alkyl group of component (2) is selected from the group consisting of methyl, ethyl, isopropyl, and t-butyl groups.
 - The resin of claim 1, wherein the vinyl aromatic component is selected from the group consisting of alpha-methylstyrene, para-methyl-alpha-methylstyrene, 2,4-diphenyl-4-methyl-1-pentene.
 - 7. A process for preparing a hydrocarbon resin of any of the preceeding claims comprising (a) providing a mixture consisting essentially of (1) about 40% to 90% by weight based on total monomers of a cyclic diolefin component comprising at least about 50% by weight dicyclopentadiene, and (2) about 60% to 10% by weight based on total monomers of at least one vinyl aromatic component having the formula

where R_1 is H or a 1-10 carbon linear or branched chain alkyl group at the meta- or para-position and R_2 is a 1-10 carbon linear or branched chain alkyl group or a 2-methyl-2-phenylpropyl group, (b) thermally polymerizing the mixture, and (c) recovering a product having a Ring and Ball softening point of about 70° to about 150°C, having a Mz with a maximum value of about 4000, having a Mw with a maximum value of about 2500, and having a Pd with a maximum value of about 3.0.

- 8. The process of claim 7, wherein the resin is subsequently hydrogenated in the presence of a catalyst comprising a metal selected from the group consisting of Group VIII, IB, VIB, and VIIB of the Periodic Table.
 - The process of claim 7, wherein low molecular weight oligomers present in the resin product are removed and are added to the mixture of components (1) and (2) for further reaction.
- 45 10. An adhesive comprising the hydrogenated resin obtainable by the process of claim 8.
 - 11. Use of the hydrocarbon resin of any one of claims 1-6 for the manufacture of caulking compounds, sealants, printing-inks, waterproofing compositions and wax compounds.
- 50 12. Use of the hydrogenated resin obtainable according to claim 8 as a tackifier for adhesives, caulking compounds, and sealants; in paints, labels, and tapes; as a modifier, extender and processing aid for plastics, printing inks, overprint varnishes and clear coatings; in textile dry sizes, ceramic tile grout, varnishes, waterproofing compositions and wax compounds.

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